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Research

Kinetics of *Xist*-induced gene silencing can be predicted from combinations of epigenetic and genomic features

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To initiate X-Chromosome inactivation (XCI), the long noncoding RNA *Xist* mediates chromosome-wide gene silencing of one X Chromosome in female mammals to equalize gene dosage between the sexes. The efficiency of gene silencing is highly variable across genes, with some genes even escaping XCI in somatic cells. A gene's susceptibility to *Xist*-mediated silencing appears to be determined by a complex interplay of epigenetic and genomic features; however, the underlying rules remain poorly understood. We have quantified chromosome-wide gene silencing kinetics at the level of the nascent transcriptome using allele-specific Precision nuclear Run-On sequencing (PRO-seq). We have developed a Random Forest machine-learning model that can predict the measured silencing dynamics based on a large set of epigenetic and genomic features and tested its predictive power experimentally. The genomic distance to the *Xist* locus, followed by gene density and distance to LINE elements, are the prime determinants of the speed of gene silencing. Moreover, we find two distinct gene classes associated with different silencing pathways: a class that requires *Xist*-repeat A for silencing, which is known to activate the SPEN pathway, and a second class in which genes are premarked by Polycomb complexes and tend to rely on the B repeat in *Xist* for silencing, known to recruit Polycomb complexes during XCI. Moreover, a series of features associated with active transcriptional elongation and chromatin 3D structure are enriched at rapidly silenced genes. Our machine-learning approach can thus uncover the complex combinatorial rules underlying gene silencing during X inactivation.

[Supplemental material is available for this article.]

X-Chromosome inactivation (XCI) is a developmental process in mammals that ensures equal gene dosage of X-linked genes between XX and XY individuals by transcriptional inactivation of one of the two X Chromosomes in female cells. In placental mammals XCI is triggered by the long noncoding RNA (lncRNA) *Xist*, which is up-regulated in a monoallelic fashion and coats the future inactive X Chromosome in *cis*, leading to the recruitment of several factors involved in transcriptional inactivation and eventually converting the entire X Chromosome into silent heterochromatin (Escamilla-Del-Arenal et al. 2011; Gendrel and Heard 2014; Galupa and Heard 2015).

Early events following *Xist* coating include the depletion of RNA polymerase II (RNAPII) from the *Xist* RNA domain and loss of active histone marks (Chaumeil et al. 2002, 2006) as well as gain of repressive chromatin modifications, such as H2AK119ub1 and H3K27me3, deposited by the Polycomb repressive complexes (PRC) 1 and 2, respectively (Plath et al. 2003, 2004; Silva et al. 2003; de Napoles et al. 2004). Subsequently, additional chromatin modifications are gained such as the histone variant macroH2A and DNA methylation of gene promoters (Escamilla-Del-Arenal et al. 2011). In recent years, progress has been made in identifying proteins that mediate *Xist*'s functions as well as the domains within the *Xist* RNA that recruit these proteins. *Xist* contains multiple conserved repeats, among which the A repeat mediates gene silencing through activation of SPEN and other factors including RBM15 (Chu et al. 2015; McHugh et al. 2015), and the B repeat recruits PRC indirectly through HNRNPK (Wutz et al. 2002; Brockdorff 2017; Pintacuda et al. 2017). The dynamics of *Xist*-mediated silencing are highly variable across genes (Chow et al. 2010; Borensztein et al. 2017; Żylicz et al. 2019), with a subset of so-called escapees remaining active even in somatic cells (Berleth et al. 2011). However, the determinants of susceptibility to XCI remain poorly understood. Because XCI is a multistep process, local interference with any step, such as *Xist* coating or access

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to the silencing machinery of one or several silencing pathways, could delay or prevent silencing of a certain gene. Defining the features that underlie differential susceptibility to XCI remains an important question, particularly because genes that are not fully silenced are implicated in diseases, such as autoimmune syndromes (Bianchi et al. 2012).

Xist RNA spreading occurs by proximity transfer to sites close to the *Xist* locus genomically or in 3D space (“*Xist* entry sites”) (Engreitz et al. 2013). From there, *Xist* has been proposed to move first into gene-dense regions and then spread to intergenic domains of the X Chromosome (Engreitz et al. 2013; Simon et al. 2013). In differentiated cells, *Xist* covers the entire X Chromosome but is reduced at escape genes (Engreitz et al. 2013; Simon et al. 2013). *Xist* coating is positively correlated with gene density and with PRC2 enrichment and negatively correlated with the density of LINE elements (Engreitz et al. 2013; Simon et al. 2013). Similarly, gene silencing tends to be slower for genes further from the *Xist* locus (Marks et al. 2015; Borensztein et al. 2017) and from LINE elements (Chow et al. 2010; Loda et al. 2017). Moreover, efficiently silenced genes tend to be enriched for Polycomb complexes (RING1B, H3K27me3) and depleted for CTCF and active marks such as H3K4me3 and H3K27ac prior to *Xist*-induced silencing (Kelsey et al. 2015; Loda et al. 2017). Thus, a variety of genetic and epigenetic features have been implicated in controlling gene-specific silencing efficiency. However, none of these features alone can predict whether and to what extent a gene will be silenced during XCI, and the associations with measured silencing efficiencies are generally weak. Because no predictive pattern of features has so far been identified, the susceptibility of genes to *Xist*-mediated silencing is likely to be controlled by a complex combination of different features.

In this study we set out to identify the genetic and chromatin features that predispose genes on the X Chromosome to be efficiently silenced or escape XCI. We measured chromosome-wide silencing dynamics of X-linked genes following induction of *Xist* expression, using allele-specific Precision nuclear Run-On sequencing (PRO-seq) (Kwak et al. 2013). We then trained two Random Forest machine-learning models to predict from 77 genomic and epigenetic features (1) whether a gene is subject to XCI, and (2) whether it will be silenced with fast or slow kinetics. Through forest-guided gene clustering, we identified feature sets that determine the silencing dynamics of subgroups of genes, indicating that variable silencing efficiencies might be associated with distinct silencing pathways. We have thus developed a framework to comprehensively assess the relative contribution of genetic and epigenetic factors to transcriptional silencing of the X Chromosome in an unbiased and quantitative manner.

Results

Quantification of gene-specific silencing dynamics by PRO-seq

To quantify gene silencing dynamics during XCI in a chromosome-wide manner, we performed allele-specific PRO-seq during ectopic *Xist* induction in female murine embryonic stem cells (mESCs). Using an inducible system allowed us to overcome the asynchronous nature of XCI, which normally limits the temporal resolution of population measurements in differentiating mESCs, the classic model system to study XCI (Chow et al. 2010). We used the female TX1072 mESC line (Schulz et al. 2014), which was derived from a cross between two different mouse strains (C57BL/6 × CAST/EiJ) and in which *Xist* up-regulation from the endogenous locus on

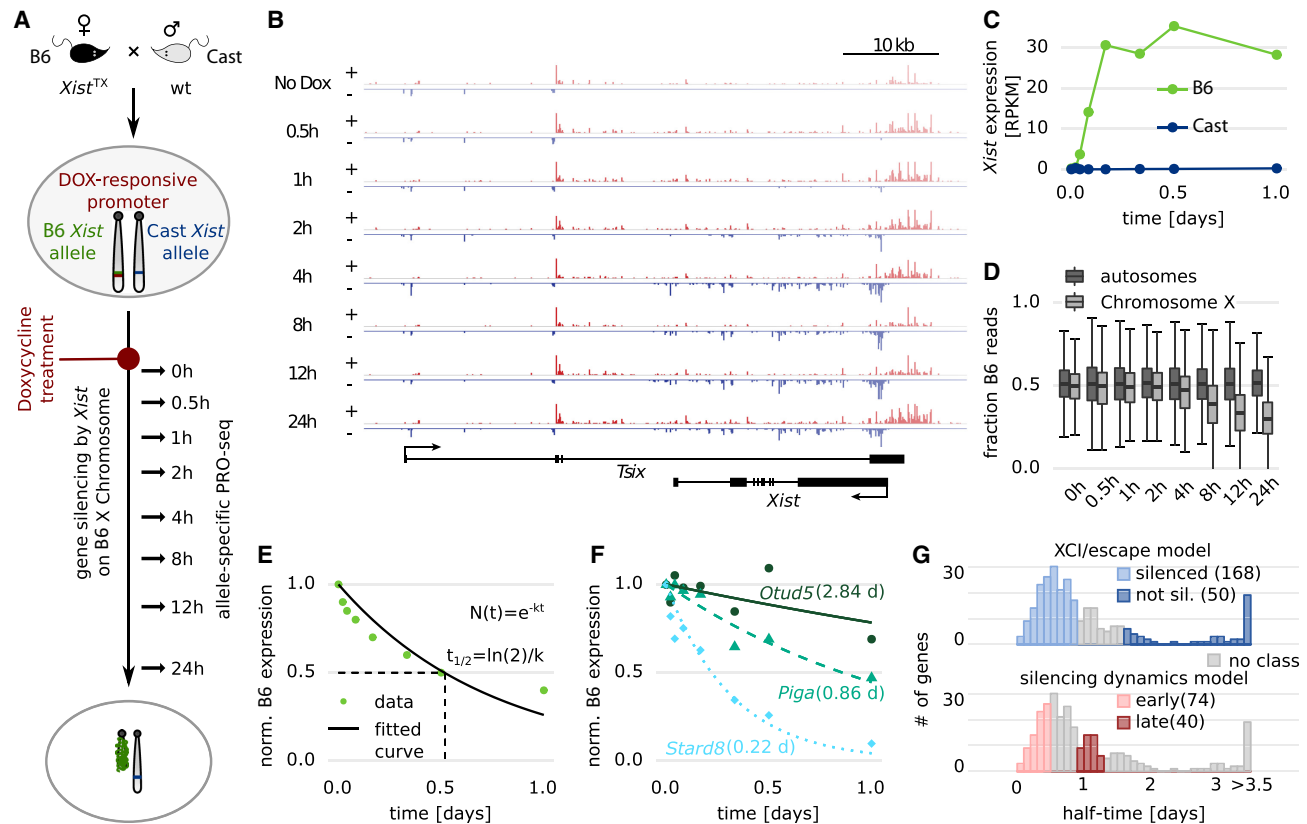
the B6 X Chromosome can be induced by doxycycline (Dox) treatment in undifferentiated cells. For a direct readout of gene silencing, we measured the nascent transcriptome by allele-specific PRO-seq (Kwak et al. 2013) at different time points up to 24 h of Dox treatment (Fig. 1A). *Xist* started to be up-regulated from the B6 chromosome about 1 h after Dox treatment and reached a plateau after 4 h (Fig. 1B,C; Supplemental Fig. S1), whereas global expression of the B6 X Chromosome was gradually reduced over time due to X inactivation, starting at 4 h of treatment (Fig. 1D).

To quantify silencing dynamics, we fitted an exponential decay function to each gene and estimated gene-specific silencing half-times, that is, the time point when transcription on the B6 X Chromosome is reduced by 50% compared to the uninduced control (Fig. 1E and examples in 1F). After several filtering steps, we had estimated reliable half-times for 280 genes, which were used for all subsequent analyses and ranged from several hours up to several days (Fig. 1G; Supplemental Tables S1, S2).

Silencing dynamics are comparable in vitro and in vivo

To ensure that the relative silencing dynamics across genes, when XCI is induced in undifferentiated mESCs, are comparable to those in the cellular context where XCI occurs endogenously, we generated two additional data sets, in which mRNA-seq was performed at different time points of Dox treatment in undifferentiated and differentiating mESCs (Fig. 2A). The computed half-times were comparable between these two data sets (Pearson correlation coefficient: $r=0.75$) (Fig. 2B), suggesting that the differentiation process only has a minor impact on relative gene silencing dynamics. When comparing half-times estimated from the two different data types (mRNA-seq vs. PRO-seq), correlation was generally a bit lower, independent of the cellular context (Pearson correlation coefficient $r=0.52$ and $r=0.51$) (Fig. 2C,D), which would be expected given that PRO-seq measures the direct transcription dynamics, whereas mRNA-seq kinetics are modulated by transcription, RNA-processing, and degradation. We also compared the estimated half-times to a previous study (Marks et al. 2015) that had used a different, Dox-independent strategy to make XCI nonrandom by inserting a stop-cassette in *Xist*'s repressive antisense transcript *Tsix*. The silencing classes defined in Marks et al. (2015) (early, intermediate, late, escapee) are in good agreement with the half-times estimated from the PRO-seq data (Fig. 2E), suggesting that Dox-induced XCI recapitulates endogenous gene silencing dynamics.

Finally, we compared our *Xist*-induced gene silencing half-times in mESCs to the dynamics of (imprinted) XCI measured in preimplantation mouse embryos in vivo through single-cell RNA-seq (Borensztein et al. 2017). The gene classification in that study (early: 16-cell stage; intermediate: 32-cell stage; late: blastocyst stage or escapee) was once more in good agreement with the silencing half-times estimated from the PRO-seq data (Fig. 2F). Based on the PRO-seq derived silencing half-times, we classified all genes according to whether they are subject to XCI or escape (Figs. 1G, 2G, silenced/not silenced) and whether they are silenced with slow or fast kinetics (Figs. 1G, 2G, early/late). Genes with intermediate half-times between the classes were excluded from the analysis (see gap between groups in Fig. 1G). The resulting classes largely agree with those previously defined in differentiating mESCs and in preimplantation embryos (compare Fig. 2G with Fig. 2E,F). Moreover, the “not silenced” class contains 37 of 50 (74%) known escapees annotated from different cell types and is strongly enriched for escapees compared to the “silenced” class when considering only (high confidence) escapees identified in



at least two different studies (e.g., *Ddx3x*, *Taf1*, *Pdcb1*, *Kdm6a*, *Usp9x*, *Hcfc1*, *Hdac6*, *Mgmt1*, *Ftx*, *Nkap*, and *Uba1*, odd ratio = 6.7, $P = 2.6 \times 10^{-5}$, Fisher's exact test) (Supplemental Table S2; Yang et al. 2010; Berletch et al. 2011; Splinter et al. 2011; Calabrese et al. 2012; Wu et al. 2014; Marks et al. 2015; Andergassen et al. 2017).

Identifying determinants of gene silencing dynamics with Random Forest modeling

We noted that genes close to the *Xist* locus tended to be silenced earlier than distal genes (Fig. 2H), in agreement with a previous study (Marks et al. 2015). However, many genes did not follow this trend. To uncover additional factors that potentially determine the susceptibility to *Xist*-mediated silencing, we developed a machine-learning model to predict silencing dynamics based on genomic and epigenetic features.

We collected 138 publicly available high-throughput data sets (ChIP-seq and bisulfite-seq) measuring chromatin modifications and DNA-binding factors, mostly in male mESCs. Because these data sets had been generated in undifferentiated mESCs, they correspond to the chromatin state before *Xist* induction. After stringent filtering on data quality, we computed the enrichment for 59 of these epigenomic features at promoters or gene body regions as appropriate (Table 1; Supplemental Text S1). In ad-

dition, we included a series of genomic and structural features, such as gene density, the frequency of 3D chromatin interactions with different genomic elements, and the linear distance to other genomic features, such as the distance to the *Xist* locus, the next TAD boundary, lamin-associated domain (LAD), or full-length LINE element (Table 1; Supplemental Text S1).

A linear model to predict a gene's susceptibility to *Xist*-mediated silencing from the collected epigenetic and genomic features had little predictive power. This indicates that no single linear combination of features or rules could be defined to discriminate, for example, silenced from not silenced genes. The different functional domains of *Xist* might recruit distinct silencing complexes (e.g., PRC1 and SPEN/HDAC3), and susceptibility to each silencing pathway might be determined by distinct feature patterns. We expected to identify such feature patterns from our data set, because genes that require the *Xist* A repeat for silencing, which activates SPEN, exhibit longer silencing half-times compared to A repeat-independent genes that might be preferentially targeted by Polycomb complexes (Kolmogorov-Smirnov [KS] test, $P = 2.2 \times 10^{-6}$) (Fig. 2I; Sakata et al. 2017).

To identify combinatorial rule sets that could predict silencing susceptibility, we used Random Forest, a nonparametric machine-learning method that combines an ensemble of single classification trees, which successively split the feature input space in a nonlinear fashion, to predict the value of a discrete binary

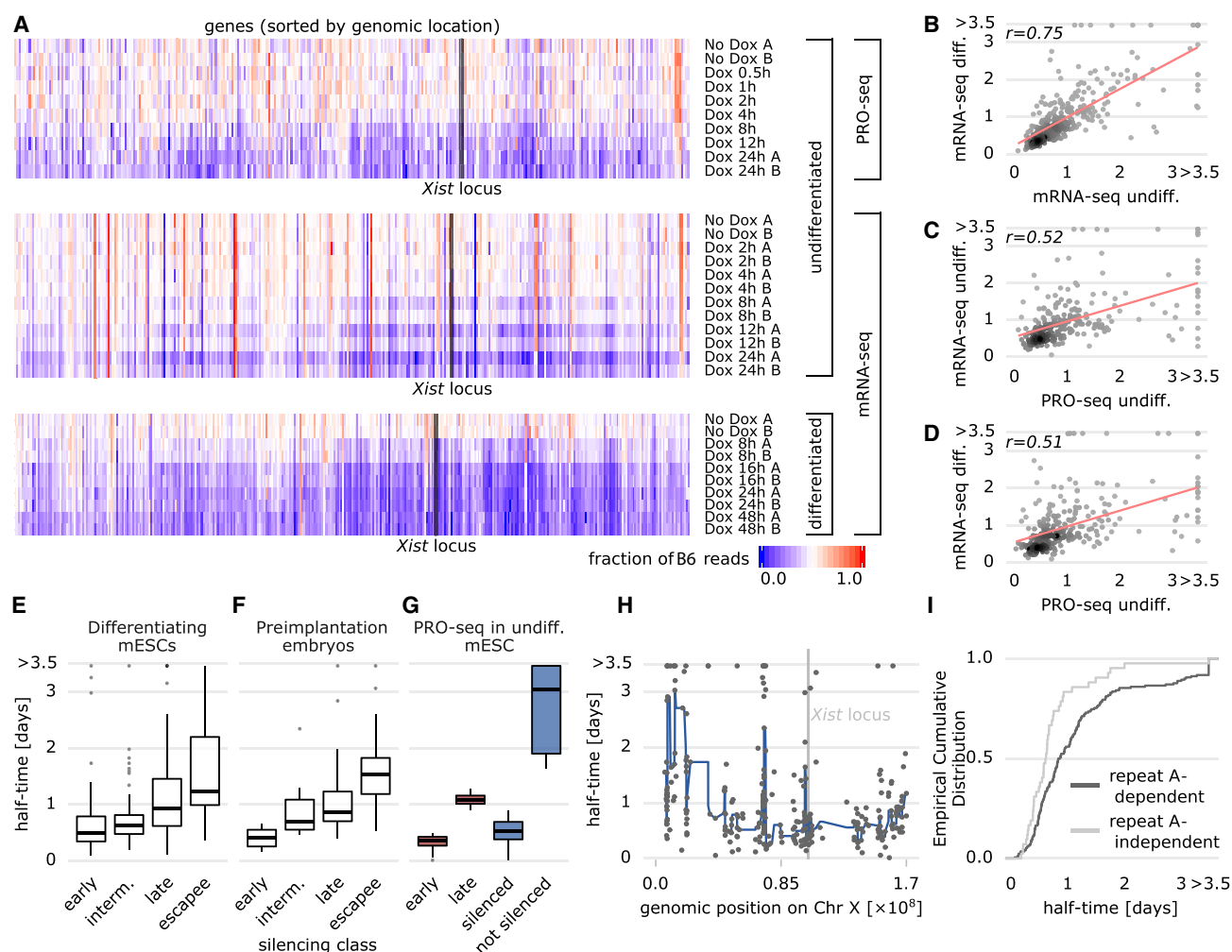


Figure 2. Comparison of PRO-seq-based silencing half-times to other data sets. (A) Comparison of PRO-seq (undifferentiated mESC, upper) and mRNA-seq data (undifferentiated mESCs, middle; differentiated mESCs, lower). Fraction of B6 reads are shown for all genes covered in all three data sets, ordered by genomic position. (B–D) Comparison of estimated half-times (in days) between the data sets shown in A (replicate B only for mRNA-seq undiff.) with fitted regression lines (red). Pearson correlation coefficients are indicated. (E–G) Distribution of half-times within silencing classes defined previously in mESCs (E) (Marks et al. 2015), in preimplantation mouse embryos (F) (Borensztein et al. 2017), and the classes used for Random Forest modeling (G): (blue) XCI/escape model; (red) silencing dynamics model. (H) Estimated half-times (black circles) for all genes in the PRO-seq data set along the X Chromosome. A fitted smooth curve of the half-times is displayed as a blue line, and the *Xist* locus is marked with a gray line. (I) Cumulative distribution of half-times of genes silenced (independent, light gray) or not silenced (dependent, dark gray) by *Xist* lacking the repeat A element (Sakata et al. 2017).

variable (Fig. 3A,B). We built two binary classification models to predict from a total of 77 epigenetic and genomic features whether a gene would be silenced or not (XCI/escape model), and whether it would be silenced early or late (silencing dynamics model) using the classification described above (Table 1; Fig. 1G). The half-time thresholds were selected such that they maximize the model classification accuracy (Supplemental Table S3; Supplemental Text S2). The XCI/escape model would identify combinations of factors, that are important for silencing in general, and the silencing dynamics model would find those that influence the kinetics of gene silencing.

The two Random Forest models predict gene silencing dynamics with error rates of 28%–29% meaning that 71%–72% of genes are classified correctly. We assessed the individual contribution of each feature to the classification accuracy via Random Forest variable importance analysis for each class in the two models by the Mean Decrease in Accuracy (MDA), which indicates the

importance of the feature for the classification performance (Fig. 4; Supplemental Fig. S2; Supplemental Text S2). We also trained our models on a set of only 10 (XCI/escape model) and 8 (silencing dynamics model) top features, which greatly improved the prediction error rate to 22.5% and 21.5%, respectively (Supplemental Fig. S3). The most important feature associated with silencing in both models was close genomic proximity of a gene's transcriptional start site (TSS) to the *Xist* locus (MDA 15%–16% in the XCI/escape model, MDA 14%–22% in the silencing dynamic model) (Fig. 4; Supplemental Figs. S2, S4, S5). Also a close proximity to LINE elements, low gene density, and enrichment for PRC1 (RING1B, H2AK119ub1, RYBP) and PRC2 (EZH2, H3K27me3) are associated with (early-) silenced genes in both models, with PRC1 playing a more prominent role in the XCI/escape model (MDA 8% for the not silenced class). (Early-) silenced genes are enriched for histone deacetylases (HDAC), involved in gene repression (MDA 1.6%–2.3% for the early-silenced class and 3.1%–

Table 1. Epigenetic and genomic features used for modeling

Epigenetic features						Genomic features	
Sequence-specific transcription factors	Structural proteins	General transcription regulators	Chromatin modifications (activation)	Chromatin modifications (repression)	Others	Genomic elements	3D structure
MYC, ESRRB, KLF4, MAFK, NANOG, MYCN, POU5F1, SOX2, TCF3, TCF3P2L1, YY1, ZNF384	CTCF, SMC1, SMC3	CDK9, E2F1, HCFC1, MAX, MED1, MED12, NIPBL, RNAPII (S2P, S5P, S7P, unphosphorylated), SIN3A, TAF1, TAF3, TBP	H3K27ac, H3K9ac, H3K4me1, H3K4me3, H3K36me3, H3K79me2, KMT2B/MLL2	H2AK119ub1, H3K27me3, RING1B (PRC1), CBX7 (PRC1) RYBP (PRC1), KMT6/EZH2 (PRC2), SUZ12 (PRC2), KDM1A/LSD1, KDM2A, KDM2B, HDAC1, HDAC2, HDAC3, DNA methylation (BS-seq), 5fC (MeDIP), 5hmC (MeDIP), TET1	H2A.Z, OGT, BRG1, CBX3	Distance to the <i>Xist</i> locus, TAD borders, LADs, full-length LINEs overlap with <i>Xist</i> entry sites, LADs, CpG islands full-length LINE density (700 kb) gene density (1 Mb) CpG content	Number Hi-C all, strength Hi-C all, number Hi-C promoter, strength Hi-C promoter, strength Hi-C <i>Xist</i> , number HiCap promoter, number HiCap enhancer, number HiCap all

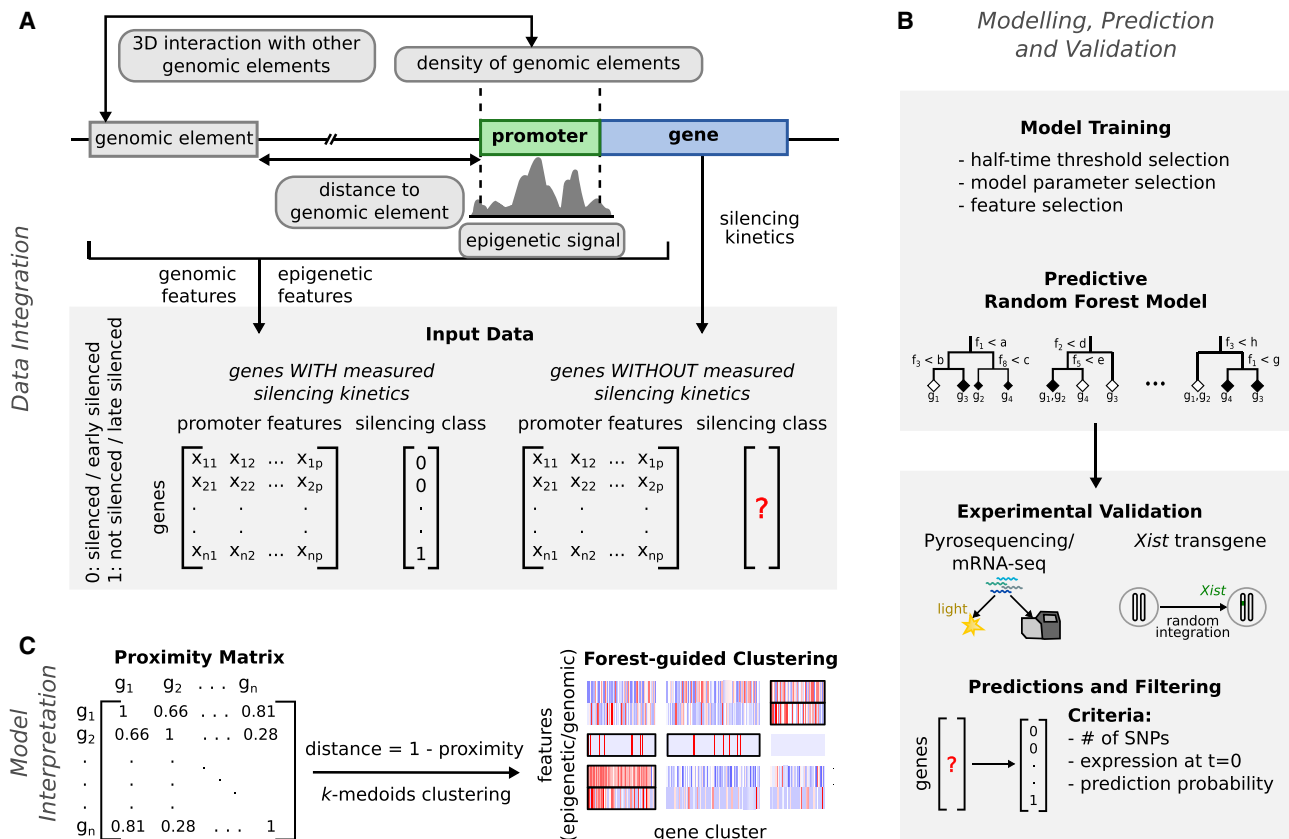
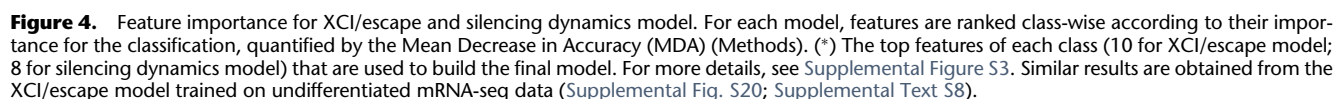
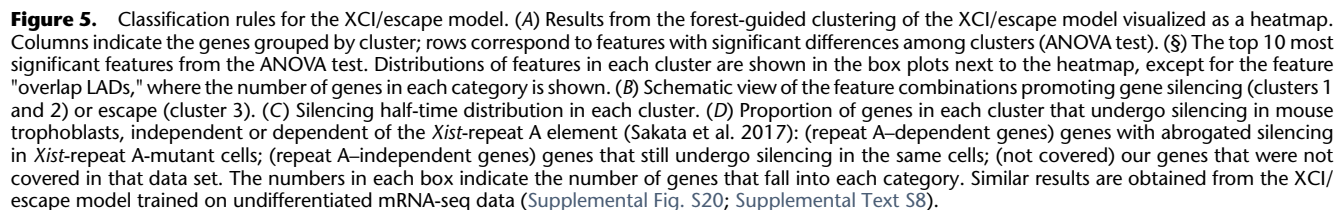


Figure 3. Schematic overview of our modeling approach. (A) Epigenetic and genomic input data for the model are collected, and feature matrices are computed for all X-linked genes with estimated half-times (labeled) and without estimated half-times (unlabeled). (B) After model training, the XCI/escape model is then used to predict the silencing class of all unlabeled X-linked genes given the same set of input features. The predictions are validated by comparing them to measured half-times from undifferentiated mRNA-seq data, with pyrosequencing experiments (few selected genes) and with measured silencing dynamics of genes in six transgenic mESCs clones. (C) A forest-guided clustering approach was developed for model interpretation. A proximity matrix between genes is computed from the trained model and converted into a distance matrix. Clusters of genes and their most significant associated features are displayed as a heatmap.



To test whether these two clusters might be associated with different silencing pathways, we analyzed how they were affected in *Xist* mutants lacking either the A or the B and C repeats (Supplemental Text S4). We used data from a previous study that had analyzed both mutations in mESCs (Bousard et al. 2018) and from another study that had characterized the A-repeat mutant in trophoblasts in vivo (Sakata et al. 2017). Cluster 2 was enriched for genes that could still be silenced by a BC-repeat mutant compared to cluster 1 (repeat BC independent, odd ratio=1.6, $P=0.19$, Fisher's exact test), whereas cluster 1 was enriched for genes still silenced in the A-repeat mutant in either data set (mESC data: odd ratio=2.3, $P=0.09$ and trophoblast data: odds ratio=2.76, $P=0.003$, Fisher's exact test) (Fig. 5D; Supplemental Fig. S9). These findings are consistent with the idea that genes that require Polycomb for silencing (repeat BC dependent) are already pre-marked by PRC1/2 and the associated histone modifications (cluster 1), in contrast to genes that require that A repeat, which activates the SPEN/HDAC3 silencing pathway (cluster 2).



analyzed the repeat A and repeat BC dependency in the two early-silenced clusters. Again, the Polycomb premarked cluster 1 tends to be enriched for repeat A-independent genes, albeit this effect was not statistically significant, and no difference was found for repeat BC dependent genes (Fig. 6D; Supplemental Fig. S9).

To validate our machine-learning model, we used the trained XCI/escape Random Forest model to predict the silencing class for X-Chromosomal genes that had not been measured in the PRO-seq experiment owing to insufficient coverage and had therefore not been used for model training (Fig. 3B; [Supplemental Table S4](#)). We performed an independent Dox induction time course experiment and used pyrosequencing to assess the allele-specific expression of six genes predicted to be silenced (Fig. 7A, top) and five genes predicted to be not silenced (Fig. 7A, bottom). The half-times

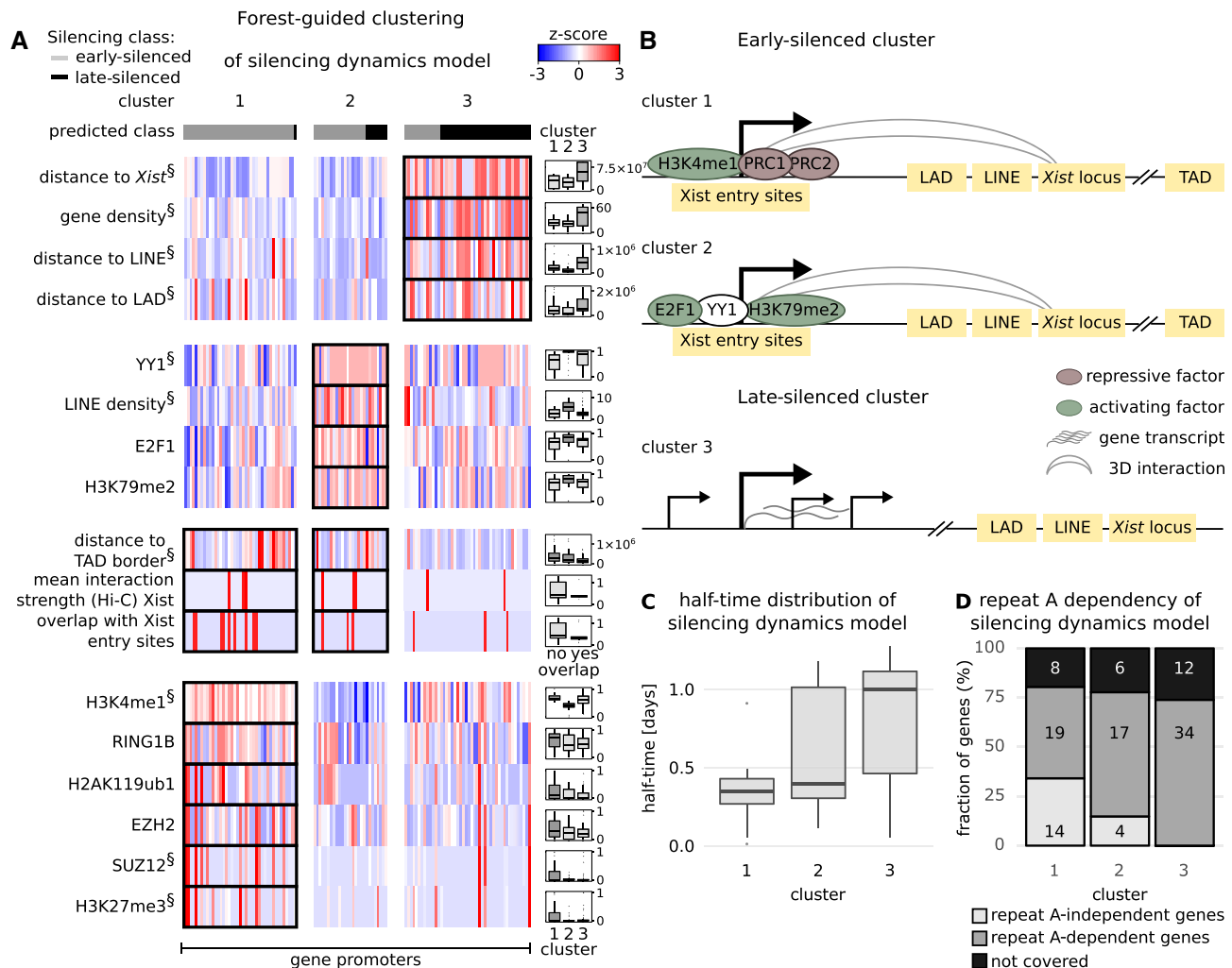


Figure 6. Classification rules for the silencing dynamics model. (A) Results from the forest-guided clustering of the silencing dynamics model visualized as a heatmap. Columns indicate the genes grouped by cluster; rows correspond to features with significant differences among clusters (ANOVA test). (§) The top 10 most significant features from the ANOVA test. Differences in the distributions of features between clusters are highlighted in the box plots next to the heatmap, except for two features, where the number of genes in each category is shown. (B) Schematic view of the features associated with early (cluster 1 and 2) and late gene silencing (cluster 3). (C) Silencing half-time distribution for each cluster. (D) The proportion of genes which undergo silencing in mouse trophoblasts, independent or dependent of the *Xist*-repeat A element, is shown for each cluster, similar to Figure 5D. The numbers in each box indicate the number of genes that fall into each category.

of the silenced genes ranged from 0.45 to 0.87 d, and those of the not silenced genes lay between 0.99 and 3.01 d. The difference was highly significant ($P=0.0043$, Wilcoxon rank-sum test) (Fig. 7B) and the half-time of all silenced genes fell in the silenced category (<0.9 d) (Fig. 1G), and three of five not silenced genes also fell in the respective category (>1.6 d) (Fig. 1G). To further validate the model, we compared all model predictions to the silencing half-times estimated from the mRNA-seq time course in undifferentiated mESCs (Fig. 2A, replicate A). Genes predicted as not silenced exhibited much longer silencing half-times than genes predicted as silenced ($P=1.7 \times 10^{-5}$, Wilcoxon rank-sum test) (Fig. 7C). Genes that have not previously been reported as escapees, but are either measured or predicted from the PRO-seq model to be not silenced and are also measured as not silenced in the mRNA-seq data, are potentially novel escape genes, such as *B630019k06Rik*, *Porcn*, *Ssr4*, *Gm14820*, and *Ppp1r3f* (Supplemental Tables S2, S4).

We next tested whether the model could predict silencing susceptibility to *Xist* transgenes located on an autosome. We used published allele-specific mRNA-seq data for a series of mESC clones that had integrated doxycycline-inducible *Xist* transgenes in different locations on the X Chromosome and on Chromosome 12 (Supplemental Table S5; Loda et al. 2017). The cells were trisomic for Chromosome 12 such that silencing of one copy should not affect cell viability. We adapted the model feature “distance to *Xist*” to account for the different locations of the transgene and calculated the fraction of genes predicted to be silenced by our model within all genes that were silenced, for each *Xist* transgene (Fig. 7D, red lines). These values varied considerably between clones depending on the size of the chromosomes and the location of the transgene. For five of six clones, the percentage predicted to be silenced was significantly higher than expected for a random sample, as estimated through a bootstrapping approach (Fig. 7D, compare red line to background distribution;

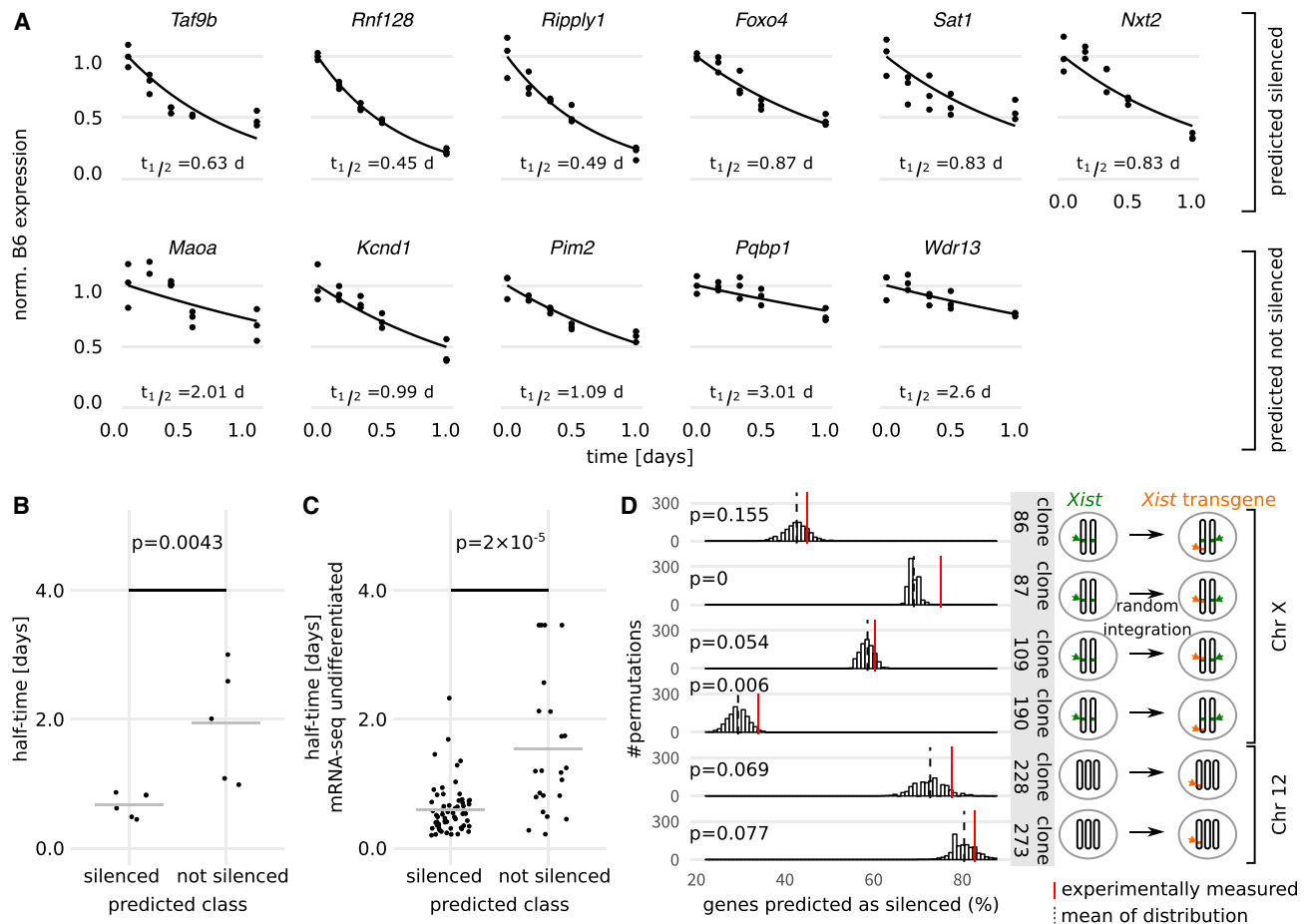


Figure 7. Experimental validation of model predictions. (A) Half-times of six candidate genes predicted as “silenced” (top) and five candidate genes predicted as “not silenced” (bottom) were estimated experimentally through allele-specific quantification by pyrosequencing at different time points during 24 h of doxycycline treatment in TX1072 cells in three independent experiments. Individual data points (dots), the fitted exponential decay function (line), and the estimated silencing half-times are shown. (B) Dot plot of the silencing half-times ($t_{1/2}$) estimated in A. (C) Dot plot of undifferentiated mRNA-seq half-times for genes predicted as silenced and not silenced by our XCI/escape model. The gray line in B and C indicates the mean, and the P -value (Wilcoxon rank-sum test) indicates a significant difference between the mean of the two distributions. (D) Fraction of genes correctly predicted as silenced by the XCI/escape model (red lines) for six cell lines in which an inducible *Xist* transgene was integrated in different chromosomal locations (orange, cartoon on the right) (Supplemental Table S5). The background distributions of silenced predictions used to estimate empirical P -values is also shown (histogram, black dashed line represents the mean).

Supplemental Text S5). Although potentially limited by the efficiency of the transgenes (Supplemental Fig. S12), this analysis shows that our model can to some extent be generalized even to other chromosomes. Taken together, these results confirm that our machine-learning model can predict X-Chromosomal gene silencing based solely on epigenetic and genomic features.

Discussion

In this study we developed a machine-learning model that can predict a gene’s susceptibility to *Xist*-mediated silencing from a combination of epigenetic and genomic features. To train the model, we measured silencing kinetics with high temporal resolution through allele-specific PRO-seq. Compared to previous studies (Marks et al. 2015; Borensztein et al. 2017), we assessed silencing dynamics by measuring nascent transcription, therefore observing instantaneous changes in transcription by transcriptionally engaged RNAPII allowing a more direct quantification of silencing compared to mRNA-seq, which was previously used. Moreover, the use of an inducible system allowed us to uncouple XCI from

differentiation and to avoid the use of mutations to ensure non-random XCI. Unlike previous studies that focused on just a few genes and/or investigated a few selected promoter features that are potentially linked to the XCI (Kelsey et al. 2015; Marks et al. 2015; Loda et al. 2017), we set out to identify silencing determinants in an unbiased manner based on a large number of epigenetic and genomic features. To uncover the combinatorial rules that control silencing dynamics, we went one step beyond classical variable importance analysis in Random Forests and introduced a random forest-guided visualization scheme. The determinants of silencing for groups of clustered genes recapitulated previous observations but also shed light on novel players or combination of features potentially controlling a gene’s susceptibility to *Xist*-mediated inactivation.

Linear distance and 3D interactions with the *Xist* locus (leading to fast *Xist* deposition at so-called “entry sites”) are thought to be the prime determinants of early *Xist* spreading (Engreitz et al. 2013). Our model found the same features to be highly predictive of gene silencing dynamics, an association that has been described previously (Marks et al. 2015; Borensztein et al. 2017) and suggests

that efficient *Xist* coating is required for fast silencing. However, previous studies had shown that *Xist* RNA initially tends to spread to gene-dense and LINE-poor regions (Engreitz et al. 2013; Simon et al. 2013), but in our analysis, gene density was associated with reduced silencing and LINE elements were found in proximity to rapidly silenced genes. A similar association was reported previously (Chow et al. 2010; Loda et al. 2017), suggesting that *Xist* coating is not the only determinant of silencing.

The *Xist* RNA recruits several protein complexes that mediate gene silencing, such as SPEN, which binds directly to the repeat A element and PRC complexes, which are indirectly recruited by the repeat B (Wutz et al. 2002; Chu et al. 2015; Monfort et al. 2015; Brockdorff 2017). Our model identified groups of genes associated with each of these silencing pathways. Repeat B/PRC-associated genes are already enriched for PRC1 and PRC2 prior to the onset of XCI, suggesting that Polycomb premarking might promote and even accelerate gene silencing and/or reinforce *Xist* spreading, as suggested by a recent study (Colognori et al. 2019). A similar enrichment of PRC components was previously found at genes susceptible to ectopic silencing by *Xist* transgenes (Kelsey et al. 2015; Loda et al. 2017). Although we did not find a clear signature at repeat A-associated silenced genes in the XCI/escape model, early-silenced genes in the silencing dynamics model that are not enriched for PRC, are located in particularly LINE-dense regions, suggesting that LINE elements might promote silencing of Polycomb-independent genes.

Previous studies looking at post-XCI cells proposed a role of CTCF in XCI (Filippova et al. 2005; Berletch et al. 2015) and found a moderate enrichment of CTCF prior to XCI at promoters of escapees compared to promoters of silenced genes (Loda et al. 2017). Although CTCF was not one of the discriminating promoter features in our XCI/escape model, we observed a significant enrichment of CTCF signal at enhancers of not silenced X-linked genes (Supplemental Fig. S13; Supplemental Text S6), suggesting a potential role of CTCF in gene silencing mediated by chromatin looping between enhancers and promoters.

Finally, our analysis identified several structural features that appear to modulate the dynamics of silencing. A high “connectivity” of some genes, that is, how much the gene is involved in 3D interactions with other genomic elements, is associated with faster silencing, maybe because *Xist* RNA can spread more easily to these genes through proximity transfer. Moreover, early silencing preferentially occurs at genes that are close to LADs, which generally contain repressed genes (van Steensel and Belmont 2017), whereas genes close to TAD boundaries tend to be silenced late.

In conclusion, we developed two Random Forest models that can accurately predict silenced and not silenced/escape genes, but also classes of early- versus late-silenced genes, constituting the first chromosome-wide predictive models of gene silencing from a very large set of features. We confirmed the predictive nature of our models by experimental testing of model predictions. The Random Forest approach allowed us to quantify the relative contribution of several features that were previously associated with XCI (e.g., linear distance to *Xist*, LINE elements, enrichment for PRC1 and PRC2, etc.) and suggested new features, which can be tested in more detail in future studies, such as TET1, and some pluripotency factors, such as ESRRB and SOX2, which have recently been implicated in reactivation of the X Chromosome during reprogramming (Janiszewski et al. 2019). Additional features could be included in the model in the future to further improve our ability to predict silencing susceptibility, and a detailed experimental investigation of the different silencing pathways elicited by *Xist* will facilitate the

interpretation of the features that predict silencing dynamics as well as escape from XCI.

Methods

ES cell culture

The female TX1072 mESC line was grown in serum-containing medium, supplemented with LIF and 2i, as previously described (Schulz et al. 2014). Details on the cell line and culture conditions are given in Supplemental Text S7.

PRO-seq

PRO-seq was performed as described previously with some modifications (Mahat et al. 2016). For details about the experiment and the allele-specific bioinformatics analysis, see Supplemental Text S7.

mRNA-seq and pyrosequencing

RNA was isolated and converted to cDNA using standard procedures. For mRNA-seq, libraries were prepared using the TruSeq Stranded mRNA LT Sample Prep Kit (Illumina) and sequenced and analyzed using standard procedures (Supplemental Text S7). For pyrosequencing, an amplicon containing a SNP is amplified by PCR from cDNA using GoTaq G2 Flexi (Promega) with 2.5 mM MgCl₂ or HotStarTaq (Qiagen) for 40 cycles. The PCR product was sequenced using the PyroMark Q24 system (Qiagen). Assay details are given in Supplemental Table S6.

Silencing half-times

To normalize for sequencing depth, the reads mapping on the B6 genome were divided by the total number of allele-specific reads for each gene as follows:

$$f_{B6}^t = \frac{reads_{B6}^t}{reads_{B6}^t + reads_{Cast}^t}. \quad (1)$$

f_{B6}^t was averaged across replicates (0, 24 h), which showed good replicate agreement (Supplemental Fig. S14), resulting in a total of eight time points ($t=0, 0.5, 1, 2, 4, 8, 12, 24$ h). The allelic ratio was calculated as follows:

$$ratio^t = \frac{f_{B6}^t}{f_{Cast}^t} = \frac{f_{B6}^t}{1 - f_{B6}^t} \quad (2)$$

and normalized to the uninduced control ($t=0$) to correct for basal skewing (different transcriptional activity at the two alleles in the absence of Dox) by the following:

$$norm^t = \frac{ratio^t}{ratio^0} = \frac{f_{B6}^t}{1 - f_{B6}^t} \times \frac{1 - f_{B6}^0}{f_{B6}^0}. \quad (3)$$

To estimate gene-specific silencing half-times, an exponential decay function was used:

$$N(t) = e^{-kt}, \quad (4)$$

where k represents the silencing rate that was fitted to $norm^t$ using the `nls` function (`stats` R package) and the half-time was calculated as

$$half-time \ t_{1/2} = \frac{\ln(2)}{k}. \quad (5)$$

A maximum value of $k=5$, corresponding to a half-time of 3.5 d was set, because higher half-times cannot be reliably estimated

from our data as a result of the limited range of time points from 0 to 24 h. The goodness of fit was evaluated via the square root of the sum of squared residuals *sqrRSS* defined as

$$\text{sqrRSS} = \sqrt{\sum_t (\text{norm}^t - N(t))^2}. \quad (6)$$

After filtering out genes without SNPs, with fewer than 10 reads for at least one time point and strong allelic skewing ($f_{B6}^0 < 0.2$ or $f_{B6}^0 > 0.8$) and *sqrRSS* > 1.5, we obtained reliable half-times for 296 X-Chromosomal genes on mouse genome mm10 (Supplemental Tables S1, S2). Those genes were mapped to the mouse genome mm9 with the *liftOver* tool from the UCSC Genome Browser (Kuhn et al. 2007).

Gene half-times from both differentiated and undifferentiated mRNA-seq time series were computed in the same way as described above. For the undifferentiated mRNA-seq data set, we discarded replicate B because of insufficient read coverage and only used replicate A (Fig. 2A), which resulted in computing half-times for 346 genes. For the differentiated mRNA-seq data set, we averaged replicate A and B for each time point and computed half-times for 379 genes. For 233 genes, half-time could be estimated from all three data sets.

Definition of model features

The epigenetic and genomic features used for the Random Forest models are summarized in Table 1 and listed in detail in Supplemental Table S7. In total, 138 ChIP-seq libraries and one bisulfite sequencing experiment on undifferentiated male mESCs (only HDAC3 was assessed in female mESCs) were collected from various sources. After performing ChIP-seq library quality control with the *deepTools* package (Ramírez et al. 2014), 58 ChIP-seq libraries and the bisulfite sequencing experiment were used in the model (Supplemental Figs. S15, S16; Supplemental Table S8; Supplemental Text S1). Epigenetic features are defined as the average ChIP-seq signal in a predefined genomic region, normalized to the signal of a matched control experiment in the same region. Read counts of each feature were normalized to the control with the R package *normR* (Supplemental Fig. S17; Helmuth et al. 2016; Kinkley et al. 2016). Although for most features the signal is extracted in a genomic region around the active gene TSS, for broader features such as elongation marks H3K36me3, RNAPII S2P, and H3K79me2, the signal was averaged over the entire gene body (Supplemental Table S7). The active TSS for each gene was identified based on the PRO-seq data at time point $t=0$ through the dREG method, which finds regions that harbor bidirectional transcription (Supplemental Fig. S18; Danko et al. 2015). In addition to epigenetic features, we defined 18 genomic features, including distance of each gene's TSS to the *Xist* locus, the next TAD border, or the next full-length LINE element; gene overlap with LADs, LINEs, and CpG islands; and strength and number of 3D chromatin interactions of gene promoters with other genomic elements. For details, see Supplemental Text S1.

Random Forest classification models

Two statistical models were developed to distinguish (1) “silenced” from “not silenced” genes, referred to as “XCI/escape model,” and (2) “early”- from “late”-silenced genes, referred to as “silencing dynamics model.” The continuous half-time values were therefore assigned to discrete classes in both models, according to fixed thresholds, which were chosen such that the error rate from the classification model (described below) would be minimized (Figs. 1G, 2G; Supplemental Table S3). Genes were defined as “silenced” for $t_{1/2} < 0.9$, as “not silenced” for $t_{1/2} > 1.6$, as “early-silenced” for $t_{1/2} < 0.5$, and as “late-silenced” for $0.9 < t_{1/2} < 1.3$. The XCI/escape

model was trained on 218 genes (168 from the “silenced” and 50 from the “not silenced” class) and the silencing dynamics model on 114 genes (74 from the “early”-silenced and 40 from the “late”-silenced class).

The two Random Forest classification models were implemented with the *randomForest* R package and use 77 predictor variables (epigenetic and genomic features), which show a variable degree of correlation between each other (Supplemental Fig. S19; R Core Team 2009). The error rate of the models is computed based on the out-of-bag (OOB) error, which is the mean prediction error over all the trees of the Random Forest. Importance of each feature is computed as “mean decrease in accuracy” (MDA) (Supplemental Text S2). Variables with large positive values of the MDA correspond to important features for the classification, whereas variables with MDA close to zero or negative correspond to noise.

Feature importance (MDA) and classification performance (OOB error) measures were further averaged over a collection of five hundred Random Forests to obtain stable results.

Simple feature selection was performed to improve the model performance by removing weaker or redundant features, which potentially introduce noise. We retained only the top 10 features from each class of the XCI/escape model and the top eight features from each class of the silencing dynamics model, which yield the model minimal error rate (Supplemental Text S2). For both models, the classification performance on the top features is reported as an average of 500 Random Forests.

Given the trained XCI/escape model, we predicted the silencing class of all X-linked genes, which were not included in the training set and chose few genes for experimental validation with pyrosequencing (Supplemental Text S2).

Forest-guided clustering for model interpretation

We can extract a *proximity matrix* from the trained Random Forest, which is a rough estimate of the distance between genes based on the proportion of times the genes are found in the same leaf node of a tree (Fig. 3C; Supplemental Text S3).

Based on this *proximity matrix*, genes were grouped into three clusters with the *k*-medoids algorithm for both models (Reynolds et al. 2006). Genes of the same silencing class (e.g., not silenced) are largely expected to cluster together according to a certain combination of epigenetic and genomic features. Given the nonlinear nature of the classification problem modeled here, we also expect, to some extent, genes from the same silencing class to be grouped in different clusters according to different combinations of features.

Similar to *k*-means clustering, *k*-medoids clustering requires setting in advance the number of clusters *k* (Supplemental Text S3). The results of the *k*-medoids clustering are visualized for both models as heatmaps, displaying the top 10 features that have a significant variation across clusters according to the *P*-value of an ANOVA test and few others which are also important for classification. Compared to classical Random Forest feature importance, the outcome of the forest-guided clustering enables an alternative interpretation of the Random Forest predictions in terms of combinatorial rules that determine the silencing state of groups of genes.

Statistical tests

Kolmogorov-Smirnov (KS) test was performed to test differences in silencing dynamics between A repeat-dependent and A repeat-independent genes. A Fisher's exact test was performed to test for enrichment of escapees in the silenced class and for A or BC repeat-dependent genes in the clusters of both models. A Wilcoxon rank-

sum test was used to test for differences in half-times between silenced and not silenced genes from the pyrosequencing experiment and for comparison with mRNA-seq data. An analysis of variance (ANOVA) test was performed to find significantly different features across clusters in both models. All statistical tests were performed in R with the *base* statistical functions package.

Data access

All raw and processed sequencing data generated in this study have been submitted to the NCBI Gene Expression Omnibus (GEO; <https://www.ncbi.nlm.nih.gov/geo/>) under accession number GSE121144. The Random Forest pipeline is available as Supplemental Code, and the scripts for all additional analysis are on GitHub (https://github.com/marsicoLab/xist_mediated_gene_silencing).

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